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INTERNACIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY POT

51 International Patent Cassificate to be:

11 International Publication Number:

WO 00/05168

C01B 3/38

A1

43 International Publication Date: Theory 1 44 1990 in

72 Inventors; and
75 Inventors/Applicants for Chacuter DYBRJAER. It [DR DR]. Non-Fronzengade 15, 3 no. 16, 2.76 Copenhagen O. DR. SEIFR CHRISTENSIN, Peter [DK DR]. Clasve, 7, 180, 164-149 Copenhagen NV (DR. LUCASSEN HANSEN Viggo [DR DR]: Bettiffevej 12, DR 27 & Brensley, DR. ROSTRET-NUELSEN, L. E. [DK DR]: Foresonej 27, DR, 2830 Vinna, DR).

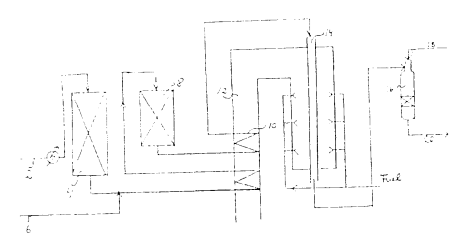
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54. Title: SYLVEHESIS GAS PRODUCTION BY STEAM REPORMING



57 Abstract

Process for the preparation of hydrogen and carbon monoxide rich, gas by steam reforming of hydrocarbon feedstock in presence Process for the preparation of myarczen and carbon monoxide rich, gas by steam reforming or myarczaroch feedstock in presence, of a steam reforming catalyst supported as that tun on the wall of a reactor, comprising steps of (a) optionally passing a puress gas. For individualist feedstock through a first react. Aith a thin film of steam reforming catalyst supported on walls of the reactor in near conducting relationship with a histogic steam, of the assing efficient from the first reactor to a subsequent tubular reactor being provided with a time nime of steam reforming catalyst and or steam reforming catalyst pellets and being neared by burning of tuel, thereby obtaining a pure sky steam reformed gas efficient and a histogram of film gas of passing the efficient trunction steams reactor to as accordance or reformer, and 3d within axing from the analytic reformer a not gas stream of product gas from in hydrogen and carron in movale.

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Synthesis Gas Production by Steam Reforming

The present invention is directed to the production of synthesis gas by steam reforming of a hydrocarbon feedstock in contact with catalyzed hardware.

The term catalyzed hardware is used for a catalyst system, where a layer of catalyst is fixed on a surface of another material, e.g. metallic surfaces. The other material serves as the supporting structure giving strength to the system. This allows to design catalyst shapes which would not have sufficient mechanical strength in itself. The system herein consists of tubes on which a thin layer of reforming catalyst is placed on the inner wall.

Alternative layouts may comprise of tubes with a catalyst layer on the outside, plates with catalyst coating, or other suitable shapes.

Synthesis gas is produced from hydrocarbons by steam reforming by the reactions (1)-(3):

$$C_n H_m + n H_2 O \rightarrow n CO + (n + \frac{m}{2}) H_2 = (-\Delta H_{298}^{O} < 0)$$
 (1)

$$CO - H_2O \rightleftharpoons CO_2 + H_2 (-\Delta H_{298}^{O} = 41 \text{ kJ/mole})$$
 (2)

$$CH_4 + H_2O = CO + 3 H_2 (-\Delta H_{298}^{\circ} = -206 \text{ kJ/mole})$$
 (3)

A second method for production of synthesis gas is autothermal reforming (ATR). In autothermal reforming, combustion of hydrocarbon feed is carried out with substoichiometric amounts of oxygen by flame reactions in a burner combustion zone and, subsequently, steam reforming of the partially combusted feedstock in a fixed bed of steam reforming catalyst. The oxidant can be air, enriched air, or pure oxygen.

A third method for production of synthesis gas is the combination of first passing the hydrocarbon feed through a fixed bed of reforming catalyst and, subsequently, passing the partly reformed feed through an autothermal reformer. The fixed bed may comprise of a number of number placed in a fired furnace. This combination is called two-step reforming or primary followed by secondary reforming and is particularly suited for production of synthesis gas for methanol and amminia production. By controlling the amount of reforming occurring in the fixed bed steam reformer before the ATR, a synthesis gas having the correct stoichiometry for methanol synthesis or a synthesis gas having the correct ratio of hydrogen to nitrogen for ammonia synthesis can be produced.

use of the art steam reforming technology makes use of reforming catalyst in the form of pellets of various sizes and shapes. The catalyst pellets are placed in fixed bed reactors (reformer tubes). The reforming reaction is endothermic. In conventional reformers, the necessary heat for the reaction is supplied from the environment outside the tubes usually by a combination of radiation and convection to the outer side of the reformer tube. The heat is transferred to the inner side of the tube by heat conduction through the tube wall and is transferred to the gas phase by convection. Finally, the heat is transferred from the gas phase to the catalyst pellet by convection. The catalyst temperature can be more than 100°C lower than the inner tube wall temperature at the same axial position of the reformer tube.

It has been found that heat transport is more efficient when catalyzed hardware is used in the steam reforming process. The heat transport to the catalyst cours by conduction from the inner tube wall. This is a much more efficient transport mechanism than the transport by convection via the gas phase. The result is that the temperatures of the inner tube wall and the catalyst are almost identical [the difference below 5°C]. Furthermore, the tube thickness can be reduced, see helow, which makes the temperature difference between the inner and outer side of the reformer tube smaller. It is hence possible to have

perature, all other conditions leng the same when replacing the occurence can refermer takes with capalyred hardware takes. A low outer take wall temperature is desirable since in prolongs the lifetime of the take. A high catalyst temperature is advantageous since the reaction rate increases with temperature and since the equilibrium of reaction (3) is shifted to the right hand side resulting in a better utilisation of the feed.

pressure drop in the datalyzed reformer tube is much lower than in the conventional case for the same tube diameter. This enables the use of reactors of non-traditional shapes e.g. tubes with small diameter and still maintaining an acceptable pressure drop. Smaller tube diameter results in an increased tube lifetime, tolerates higher temperatures and reduces the tube material consumption.

Finally, the catalyst amount is reduced when using catalyzed hardware reformer tubes compared to the conventional reformer with a fixed bed of reforming catalyst.

The small amount of catalyst distates the use of a feedstock free of catalyst poisons. This can e.g. be obtained by sending the feedstock through a prereformer.

Fig. 1 shows an example of a plant producing sympass. Feed 2 is preheated, desulphurized in unit 4, mixed with process steam 6, and further heated before entering an adiabatic prerefermer 8. The effluent stream from prerefermer 8 is further heated in a heat exchanger coil arranged in flue gas channel 12 and send to the tubular refermer 14, where conversion of methane to hydrogen, carbon monoxide, and carbon dickide occurs. The effluent gas is passed to autothermal refermer 16, wherein combustion is performed with exidant stream 18. The processing of effluent gas 20 downstream from the autothermal refermer depends on the intended use of the product.

Catalyzed nardward can be used in two of the units elown in Fig. 1:

- in the preheater coul 10 for heating the prereformer efficient gas before entering the number reformer 14.
- 2. In the tubular reformer 14.

This invention provides process for the preparation of hydrogen and carbon monoxide rich gas by steam reforming of a hydrocarbon feedstock in presence of a steam reforming catalyst supported as thin film on the wall of a reactor, comprising steps of

- (a) optionally passing a process gas of hydrocarbon feedstock through a first reactor with a thin film of steam reforming catalyst supported on walls of the reactor in heat conducting relationship with a hot gas stream;
- (b) passing effluent from the first reactor to a subsequent tubular reactor being provided with a thin film of steam reforming catalyst and/or steam reforming catalyst pellets and being heated by burning of fuel, thereby obtaining a partially steam reformed gas effluent and a hot gas stream of flue gas;
- (σ) passing the effluent from the second reactor to an autothermal reformer; and
- (d) withdrawing from the autithermal reformer a hot gas stream of product gas rich in hydrogen and carbon monoxide.

Example 1

A cutalyzed hardware refirmer reactive has been togotod. The test unit conclusts of a system for providing the feeds to the reactor, the reactor otpelf, and equipment for postprestment and analysis of the effluent gas from the reactor.

The reactor consists of a 1/4" tupe of length 1080 mm which is. In the middle 500 mm. stated on the inner wall with RMMR nickel steam reforming catalyst. The catalyst has the same composition as the REME pelletshaped steam reformind datalyst available from Halder Tepsoe A.S. The thickness of the catalyst layer is 0.31 mm. The catalyzed readtor tube is placed in a casing made of solid metal, which has a hole plosing tightly around the datalyzed tube. A number of milled grooves, in which thermoscuples are placed, is made along the hole. One of the thermocouples is movable so that a wall temperature profile of the catalyzed tube can be obtained. Additionally, thermocouples are placed in the gas channel in the catalyzed tube measuring the gas temperature at the inlet to and the outlet from the catalyzed cone. The reactor with casing is placed in an electrically heated oven, in which the temperature can be controlled seperately in 6 different cones.

The feed steams consist of hydrogen, methane, carbondickide, and steam. The feed streams are mixed and preheated before entering the reactor. After the reactor, the effluent gas stream is couled down, the condensed water is separated from the gas, and the gas composition is measured by a gaschromatograph.

Two sets of conditions were tested. One set at lower temperature to simulate use of catalyzed hardware in a preheater coil (test No. 1), and one set at higher temperature to simulate a tubular reformer "test No. 2). The conditions are shown in Table 1. The pressure was in both cases 18 bar g. The temperature profile imposed on the

reactor wall measured by the movable thermocouple is shown in Fig. 2.

Table 1

	Conditio	ons for Ca	talyzed Har	rdware Read	tor Test	
Test No.	Gas tempe- rature at inlet of cat- alyzed zone	Gas tempe- rature at out- let of catal- yzed zone	Hydro- gen flow rate	Meshane flow rate	Carbon- dioxide flow rate	Steam Elow rate
	°C	°C	N1/h	N1/h	N1/h	Nl.h
1	605	633	62.0	310.0	16.1	791.4
2	679	795	240.5	152.0	63.1	425.0

The measured effluent gas composition is shown in Table 2. The gas composition is on dry basis.

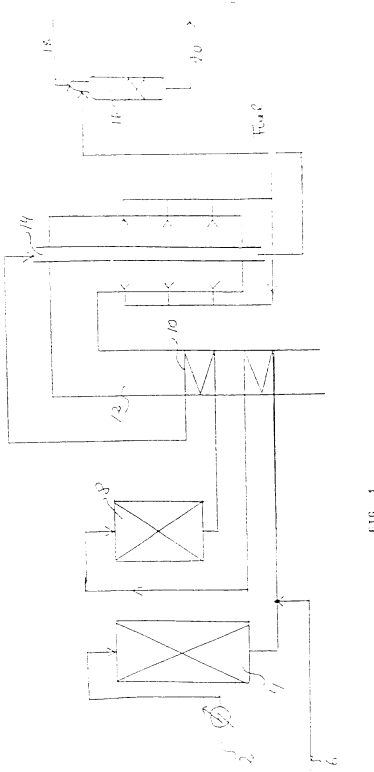
Table 2

	Effluent Gas	3 Composition c	on Dry Basis	
Test No.	Hydrogen	Carbon- monoxide	Carbon- dioxide	Methane
	mole%	mole%	molet	moles
1	49.6	2.48	11.4	36.5
2	67.8	10.8	9.80	11.7

The effluent gas is in both cases in equilibrium with respect to the reforming reaction at the outlet gas temperature within experimental uncertainty. This demonstrates that a conversion similar to a fixed bed reactor can be obtained in a catalyzed hardware reactor.

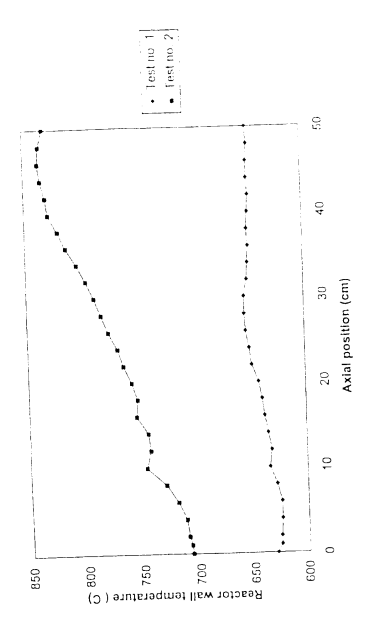
CLAIMS

- process for the preparation of hydrogen and carbon monoxide rich gas by steam reforming of a hydrocarbon feedstock in presence of a steam reforming catalyst supported as thin film on the wall of a reactor, comprising steps of
- (a) optionally passing a process gas of hydrocarbon feedstock through a first reactor with a thin film of steam reforming catalyst supported on walls of the reactor in heat conducting relationship with a hot gas stream;
- (b) passing effluent from the first reactor to a subsequent tubular reactor being provided with a thin film of steam reforming catalyst and/or steam reforming catalyst pellets and being heated by burning of fuel, thereby obtaining a partially steam reformed gas effluent and a hot gas stream of flue gas;
- (c) passing the effluent from the second reactor to an autothermal reformer; and
- (d) withdrawing from the autothermal reformer a hot gas stream of product gas rich in hydrogen and carbon monoxide.
- 2. Process of claim 1, wherein the steam reforming catalyst comprises nickel and/or noble metals.



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FIG. 2



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